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Publication Title:

PYRROLO[2,3-d]PYRIMIDINE AND THEIR USE AS PURINERGIC RECEPTOR ANTAGONISTS

Abstract:

Abstract of WO02055084

Use of a compound of the formula (I) wherein R1 is selected from alkyl, alkoxy, aryloxy, alkylthio, arylthio, aryl, halogen, CN, NR7R8, NR6COR7, NR6CO2R9 and NR6SO2R9; R2 is selected from aryl attached via an unsaturated carbon; R3 and R4 are independently selected from H, alkyl, halogen, alkoxy, alkylthio, CN and NR7R8; R5 is sel 11f1 ected from H, acyclic alkyl, COR6, CONR7R8, CONR6NR7R8, CO2R9 and SO2R9; R6, R7 and R8 are independently selected from H, alkyl and aryl or where R7 and R8 are in an NR7R8 group R7 and R8 may be linked to form a hererocyclic group, or where R6, R7 and R8 are in a (CONR6NR7R8) group, R6 and R7 may be linked to form a hererocyclic group; and R9 is selected from alkyl and aryl, or a pharmaceutically acceptable salt thereof or prodrug thereof, in the manufacture of a medicament for the treatment or prevention of a disorder in which the blocking of purine receptors, particularly adenosine receptors and more particularly A2A receptors, may be beneficial, particularly wherein said disorder is a movement disorder such as Parkinson's disease or said disorder is depression, cognitive or memory impairment, acute or chronic pain, ADHD or narcolepsy, or for neuroprotection in a subect; compounds of formula (I) for use in therapy; and novel compounds of formula (I) per se. Data supplied from the esp@cenet database - Worldwide

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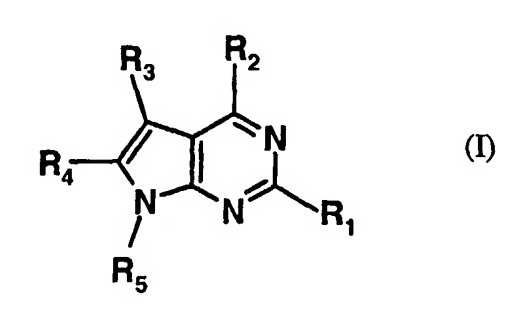
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(54) Title: PYRROLO[2,3-d]PYRIMIDINE AND THEIR USE AS PURINERGIC RECEPTOR ANTAGONISTS



(57) Abstract: Use of a compound of the formula (I) wherein R₁ is selected from alkyl, alkoxy, aryloxy, alkylthio, arylthio, aryl, halogen, CN, NR₇R₈, NR₆COR₇, NR₆CO₂R₉ and NR₆SO₂R₉; R₂ is selected from aryl attached via an unsaturated carbon; R₃ and R₄ are independently selected from H, alkyl, halogen, alkoxy, alkylthio, CN and NR₇R₈; R₅ is selected from H, acyclic alkyl, COR₆, CONR₇R₈, CONR₆NR₇R₈, CO₂R₉ and SO₂R₉; R₆, R₇ and R₈ are independently selected from H, alkyl and aryl or where R₇ and R₈ are in an NR₇R₈ group R₇ and R₈ may be linked to form a hererocyclic group, or where R₆, R₇ and R₈ are in a (CONR₆NR₇R₈) group, R₆ and R₇ may be linked to form a hererocyclic group; and R₉ is selected from alkyl and aryl, or a pharmaceutically acceptable salt thereof or prodrug thereof, in the

manufacture of a medicament for the treatment or prevention of a disorder in which the blocking of purine receptors, particularly adenosine receptors and more particularly A_{2A} receptors, may be beneficial, particularly wherein said disorder is a movement disorder such as Parkinson's disease or said disorder is depression, cognitive or memory impairment, acute or chronic pain, ADHD or narcolepsy, or for neuroprotection in a subect; compounds of formula (I) for use in therapy; and novel compounds of formula (I) per se.



PYRROLO[2,3-d]PYRIMIDINE AND THEIR USE AS PURINERGIC RECEPTOR ANTAGONISTS

The present invention relates to pyrrolo[2,3-d]pyrimidine derivatives and their use in therapy. In particular, the present invention relates to the treatment of disorders in which the reduction of purinergic neurotransmission could be beneficial. The invention relates in particular to blockade of adenosine receptors and particularly adenosine A_{2A} receptors, and to the treatment of movement disorders such as Parkinson's disease.

Movement disorders constitute a serious health problem, especially amongst the elderly sector of the population. These movement disorders are often the result of brain lesions. Disorders involving the basal ganglia which result in movement disorders include Parkinson's disease, Huntington's chorea and Wilson's disease. Furthermore, dyskinesias often arise as sequelae of cerebral ischaemia and other neurological disorders.

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There are four classic symptoms of Parkinson's disease: tremor, rigidity, akinesia and postural changes. The disease is also commonly associated with depression, dementia and overall cognitive decline. Parkinson's disease has a prevalence of 1 per 1,000 of the total population. The incidence increases to 1 per 100 for those aged over 60 years. Degeneration of dopaminergic neurones in the substantia nigra and the subsequent reductions in interstitial concentrations of dopamine in the striatum are critical to the development of Parkinson's disease. Some 80% of cells from the substantia nigra need to be destroyed before the clinical symptoms of Parkinson's disease are manifested.

Current strategies for the treatment of Parkinson's disease are based on transmitter replacement therapy (L-dihydroxyphenylacetic acid (L-DOPA)), inhibition of monoamine oxidase (e.g. Deprenyl®), dopamine receptor agonists (e.g. bromocriptine and apomorphine) and anticholinergics (e.g. benztrophine, orphenadrine). Transmitter replacement therapy in particular does not provide consistent clinical benefit, especially after prolonged treatment when "on-off" symptoms develop, and this treatment has also been associated with involuntary movements of athetosis and chorea, nausea and vomiting. Additionally current therapies do not treat the underlying neurodegenerative disorder resulting in a continuing cognitive decline in patients. Despite new drug approvals, there is

still a medical need in terms of improved therapies for movement disorders, especially Parkinson's disease. In particular, effective treatments requiring less frequent dosing, effective treatments which are associated with less severe side-effects, and effective treatments which control or reverse the underlying neurodegenerative disorder, are required.

Blockade of A₂ adaposine receptors has recently been implicated in the treatment of movement disorders such as Parkinson's disease (Richardson, P.J. et al., Trends Pharmacol. Sci. 1997, 18, 338-344) and in the treatment of cerebral ischaemia (Gao, Y. and Phillis, J.W., Life Sci. 1994, 55, 61-65). The potential utility of adenosine A_{2A} receptor antagonists in the treatment of movement disorders such as Parkinson's Disease has recently been reviewed (Mally, J. and Stone, T.W., CNS Drugs, 1998, 10, 311-320).

Adenosine is a naturally occurring purine nucleoside which has a wide variety of well15 documented regulatory functions and physiological effects. The central nervous system
(CNS) effects of this endogenous nucleoside have attracted particular attention in drug
discovery, owing to the therapeutic potential of purinergic agents in CNS disorders
(Jacobson, K.A. et al., J. Med. Chem. 1992, 35, 407-422). This therapeutic potential has
resulted in considerable recent research endeavour within the field of adenosine receptor
20 agonists and antagonists (Bhagwhat, S.S.; Williams, M. Exp. Opin. Ther. Patents 1995,
5,547-558).

Adenosine receptors represent a subclass (P₁) of the group of purine nucleotide and nucleoside receptors known as purinoreceptors. The main pharmacologically distinct adenosine receptor subtypes are known as A₁, A_{2A}, A_{2B} (of high and low affinity) and A₃ (Fredholm, B.B., *et al.*, *Pharmacol. Rev.* 1994, 46, 143-156). The adenosine receptors are present in the CNS (Fredholm, B.B., *News Physiol. Sci.*, 1995, 10, 122-128).

The design of P₁ receptor-mediated agents has been reviewed (Jacobson, K.A., Suzuki, F., Drug Dev. Res., 1997, 39, 289-300; Baraldi, P.G. et al., Curr. Med. Chem. 1995, 2, 707-722), and such compounds are claimed to be useful in the treatment of cerebral ischemia or neurodegenerative disorders, such as Parkinson's disease (Williams, M. and Burnstock, G.

Purinergic Approaches Exp. Ther. (1997), 3-26. Editor: Jacobson, Kenneth A.; Jarvis, Michael F. Publisher: Wiley-Liss, New York, N.Y.)

It has been speculated that xanthine derivatives such as caffeine may offer a form of treatment for attention-deficit hyperactivity disorder (ADHD). A number of studies have demonstrated a beneficial effect of caffeine on controlling the symptoms of ADHD (Garfinkel, B.D. et al., Psychiatry, 1981, 26, 395-401). Antagonism of adenosine receptors is thought to account for the majority of the behavioural effects of caffeine in humans and thus blockade of adenosine A_{2A} receptors may account for the observed effects of caffeine in ADHD patients. Therefore a selective A_{2A} receptor antagonist may provide an effective treatment for ADHD but without the unwanted side-effects associated with current therapy.

Adenosine receptors have been recognised to play an important role in regulation of sleep patterns, and indeed adenosine antagonists such as caffeine exert potent stimulant effects and can be used to prolong wakefulness (Porkka-Heiskanen, T. et al., Science, 1997, 276, 1265-1268). Recent evidence suggests that a substantial part of the actions of adenosine in regulating sleep is mediated through the adenosine A_{2A} receptor (Satoh, S., et al., Proc. Natl. Acad. Sci., USA, 1996). Thus, a selective A_{2A} receptor antagonist may be of benefit in counteracting excessive sleepiness in sleep disorders such as hypersomnia or narcolepsy.

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It has recently been observed that patients with major depression demonstrate a blunted response to adenosine agonist-induced stimulation in platelets, suggesting that a dysregulation of A_{2A} receptor function may occur during depression (Berk, M. et al, 2001, Eur. Neuropsychopharmacol. 11, 183-186). Experimental evidence in animal models has shown that blockade of A_{2A} receptor function confers antidepressant activity (El Yacoubi, M et al. Br. J. Pharmacol. 2001, 134, 68-77). Thus, A_{2A} receptor antagonists may offer a novel therapy for the treatment of major depression and other affective disorders in patients.

The pharmacology of adenosine A_{2A} receptors has been reviewed (Ongini, E.; Fredholm, B.B. Trends Pharmacol. Sci. 1996, 17(10), 364-372). One potential underlying mechanism in the aforementioned treatment of movement disorders by the blockade of A_2 adenosine receptors is the evidence of a functional link between adenosine A_{2A} receptors to dopamine

D₂ receptors in the CNS. Some of the early studies (e.g. Ferre, S. et al., Stimulation of high-affinity adenosine A₂ receptors decreases the affinity of dopamine D₂ receptors in rat striatal membranes. Proc. Natl. Acad. Sci. U.S.A. 1991, 88, 7238-41) have been summarised in two more recent articles (Fuxe, K. et al., Adenosine Adenine Nucleotides Mol. Biol. Integr. Physiol., [Proc. Int. Symp.], 5th (1995), 499-507. Editors: Belardinelli, Luiz; Pelleg, Amir. Publisher: Kluwer, Boston, Mass.; Ferre, S. et al., Trends Neurosci. 1997, 20, 482-487).

As a result of these investigations into the functional role of adenosine A_{2A} receptors in the CNS, especially *in vivo* studies linking A₂ receptors with catalepsy (Ferre *et al.*, *Neurosci. Lett.* 1991, 130, 162-4; Mandhane, S.N. *et al.*, *Eur. J. Pharmacol.* 1997, 328, 135-141) investigations have been made into agents which selectively bind to adenosine A_{2A} receptors as potentially effective treatments for Parkinson's disease.

- While many of the potential drugs for treatment of Parkinson's disease have shown benefit in the treatment of movement disorders, an advantage of adenosine A_{2A} antagonist therapy is that the underlying neurodegenerative disorder may also be treated. The neuroprotective effect of adenosine A_{2A} antagonists has been reviewed (Ongini, E.; Adami, M.; Ferri, C.; Bertorelli, R., Ann. N. Y. Acad. Sci. 1997, 825(Neuroprotective Agents), 30-48). In particular, compelling recent evidence suggests that blockade of A_{2A} receptor function confers neuroprotection against MPTP-induced neurotoxicity in mice (Chen, J-F., J. Neurosci. 2001, 21, RC143). In addition, several recent studies have shown that consumption of dietary caffeine, a known adenosine A_{2A} receptor antagonist, is associated with a reduced risk of Parkinson's disease in man (Ascherio, A. et al, Ann Neurol., 2001, 50, 56-63; Ross G W, et al., JAMA, 2000, 283, 2674-9). Thus, A_{2A} receptor antagonists may offer a novel treatment for conferring neuroprotection in neurodegenerative diseases such as Parkinson's disease.
- Xanthine derivatives have been disclosed as adenosine A_2 receptor antagonists as useful for treating various diseases caused by hyperfunctioning of adenosine A_2 receptors, such as Parkinson's disease (see, for example, EP-A-565377).

One prominent xanthine-derived adenosine A_{2A} selective antagonist is CSC [8-(3-chlorostyryl)caffeine] (Jacobson *et al.*, *FEBS Lett.*, 1993, 323, 141-144).

Theophylline (1,3-dimethylxanthine), a bronchodilator drug which is a mixed antagonist at adenosine A₁ and A_{2A} receptors, has been studied clinically. To determine whether a formulation of this adenosine receptor antagonist would be of value in Parkinson's disease an open trial was conducted on 15 Parkinsonian patients, treated for up to 12 weeks with a slow release oral theophylline preparation (150 mg/day), yielding serum theophylline levels of 4.44 mg/L after one week. The patients exhibited significant improvements in mean objective disability scores and 11 reported moderate or marked subjective improvement (Mally, J., Stone, T.W. *J. Pharm. Pharmacol.* 1994, 46, 515-517).

KF 17837 [(E)-8-(3,4-dimethoxystyryl)-1,3-dipropyl-7-methylxanthine] is a selective adenosine A_{2A} receptor antagonist which on oral administration significantly ameliorated the cataleptic responses induced by intracerebroventricular administration of an adenosine A_{2A} receptor agonist, CGS 21680. KF 17837 also reduced the catalepsy induced by haloperidol and reserpine. Moreover, KF 17837 potentiated the anticataleptic effects of a subthreshold dose of L-DOPA plus benserazide, suggesting that KF 17837 is a centrally active adenosine A_{2A} receptor antagonist and that the dopaminergic function of the nigrostriatal pathway is potentiated by adenosine A_{2A} receptor antagonists (Kanda, T. et al., Eur. J. Pharmacol. 1994, 256, 263-268). The structure activity relationship (SAR) of KF 17837 has been published (Shimada, J. et al., Bioorg. Med. Chem. Lett. 1997, 7, 2349-2352). Recent data has also been provided on the A_{2A} receptor antagonist KW-6002 (Kuwana, Y et al., Soc. Neurosci. Abstr. 1997, 23, 119.14; and Kanda, T. et al., Ann. Neurol. 1998, 43(4), 507-513).

New non-xanthine structures sharing these pharmacological properties include SCH 58261 and its derivatives (Baraldi, P.G. et al., Pyrazolo[4,3-e]-1,2,4-triazolo[1,5-c]pyrimidine Derivatives: Potent and Selective A_{2A} Adenosine Antagonists. J. Med. Chem. 1996, 39, 1164-71). SCH 58261 (7-(2-phenylethyl)-5-amino-2-(2-furyl)-pyrazolo-[4,3-e]-1,2,4-triazolo[1,5-c] pyrimidine) is reported as effective in the treatment of movement disorders (Ongini, E. Drug Dev. Res. 1997, 42(2), 63-70) and has been followed up by a later series of compounds (Baraldi, P.G. et al., J. Med. Chem. 1998, 41(12), 2126-2133).

The foregoing discussion indicates that a potentially effective treatment for movement disorders in humans would comprise agents which act as antagonists at adenosine A_{2A} receptors.

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It has now been found that pyrrolo[2,3-d]pyrimidine derivatives, which are structurally unrelated to known adenosine receptor antagonists, exhibit unexpected antagonist binding affinity at adenosine (P₁) receptors, and in particular at the adenosine A_{2A} receptor. Such compounds may therefore be useful for the treatment of disorders in which the blocking of purine receptors, particularly adenosine receptors and more particularly adenosine A_{2A} receptors, may be beneficial. In particular such compounds may be suitable for the treatment of movement disorders, such as disorders of the basal ganglia which result in dyskinesias. Disorders of particular interest in the present invention include Parkinson's disease, Alzheimer's disease, spasticity, Huntington's chorea and Wilson's disease.

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Such compounds may also be particularly suitable for the treatment of depression, cognitive or memory impairment including Alzheimer's disease, acute or chronic pain, ADHD, narcolepsy or for neuroprotection.

According to the present invention there is provided the use of a compound of formula (I):

$$R_4$$
 R_5
 R_5
 R_5

5 wherein

R₁ is selected from alkyl, alkoxy, aryloxy, alkylthio, arylthio, aryl, halogen, CN, NR₇R₈, NR₆CO₈, NR₆CO₈, NR₆CO₈, NR₆CO₈, and NR₆SO₈;

R₂ is selected from aryl attached via an unsaturated carbon;

R₃ and R₄ are independently selected from H, alkyl, halogen, alkoxy, alkylthio, CN and NR₇R₈;

R₅ is selected from H, acyclic alkyl, COR₆, CONR₇R₈, CONR₆NR₇R₈, CO₂R₉ and SO₂R₉; R₆, R₇ and R₈ are independently selected from H, alkyl and aryl or where R₇ and R₈ are in an NR₇R₈ group, R₇ and R₈ may be linked to form a heterocyclic group, or where R₆, R₇ and R₈ are in a (CONR₆NR₇R₈) group, R₆ and R₇ may be linked to form a heterocyclic group; and

15 R₉ is selected from alkyl and aryl,

or a pharmaceutically acceptable salt or prodrug thereof, in the manufacture of a medicament for the treatment or prevention of a disorder in which the blocking of purine receptors, particularly adenosine receptors and more particularly A_{2A} receptors, may be beneficial.

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As used herein the term "alkyl", unless otherwise stated, means a branched or unbranched, cyclic or acyclic, saturated or unsaturated (e.g. alkenyl or alkynyl) hydrocarbyl radical which may be substituted or unsubstituted. Where cyclic, the alkyl group is preferably C₃ to C₁₂, more preferably C₅ to C₁₀, more preferably C₅, C₆ or C₇. Where acyclic, the alkyl group is preferably C₁ to C₁₀, more preferably C₁ to C₆, more preferably methyl, ethyl, propyl (n-propyl or isopropyl), butyl (n-butyl, isobutyl or tertiary-butyl) or pentyl (including n-pentyl and isopentyl), more preferably methyl. It will be appreciated therefore that the term "alkyl" as used herein, unless otherwise stated, includes alkyl (branched or unbranched), alkenyl (branched or unbranched), alkenyl (branched or unbranched), cycloalkynyl.

As used herein, the term "lower alkyl" means methyl, ethyl, propyl (n-propyl or isopropyl) or butyl (n-butyl, isobutyl or tertiary-butyl).

As used herein, the term "aryl" means an aromatic group, such as phenyl or naphthyl (preferably phenyl), or a heteroaromatic group containing one or more heteroatom(s) preferably selected from N, O and S, such as pyridyl, pyrrolyl, quinolinyl, furanyl, thienyl, oxadiazolyl, thiadiazolyl, thiazolyl, oxazolyl, isoxazolyl, pyrazolyl, triazolyl, imidazolyl, pyrimidinyl, indolyl, pyrazinyl or indazolyl.

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As used herein, the term "heteroaryl" means an aromatic group containing one or more heteroatom(s) preferably selected from N, O and S, such as pyridyl, pyrrolyl, quinolinyl, furanyl, thienyl, oxadiazolyl, thiadiazolyl, thiazolyl, oxazolyl, isoxazolyl, pyrazolyl, triazolyl, imidazolyl, pyrimidinyl, indolyl, pyrazinyl or indazolyl.

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As used herein, the term "non-aromatic heterocyclyl" means a non-aromatic cyclic group containing one or more heteroatom(s) preferably selected from N, O and S, such as a cyclic amino group (including aziridinyl, azetidinyl, pyrrolidinyl, piperidyl, piperazinyl, morpholinyl) or a cyclic ether (including tetrahydrofuranyl).

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As used herein, the term "alkoxy" means alkyl-O-. As used herein, the term "aryloxy" means aryl-O-.

As used herein, the term "halogen" means a fluorine, chlorine, bromine or iodine radical.

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As used herein, the term "ortho, ortho-disubstituted aryl groups" refers to aryl groups which are substituted in both ortho positions of the aryl group relative to the point of attachment of the aryl group to the pyrimidine ring.

30 As used herein, the term "prodrug" means any pharmaceutically acceptable prodrug of a compound of the present invention.

Where any of R₁ to R₁₅ is selected from alkyl, alkoxy and thioalkyl, in accordance with formula (I) as defined above, then that alkyl group, or the alkyl group of the alkoxy or thioalkyl group, may be substituted or unsubstituted. Where any of R₁ to R₁₅ are selected from aryl, aryloxy and thioaryl, in accordance with formula (I) as defined above, then said aryl group, or the aryl group of the aryloxy or thioaryl group, may be substituted or unsubstituted. Where R₇ and R₈, or R₆ and R₇, are linked to form a heterocyclic group, the heterocyclic group may be substituted or unsubstituted. Where substituted, there will generally be 1 to 3 substituents present, preferably 1 substituents. Substituents may include: carbon-containing groups such as

10 alkyl,

aryl, (e.g. substituted and unsubstituted phenyl (including

(alkyl)phenyl, (alkoxy)phenyl, (alkyl- and aryl-

sulfonylamino)phenyl and halophenyl),

arylalkyl; (e.g. substituted and unsubstituted benzyl);

15 halogen atoms and halogen containing groups such as

haloalkyl (e.g. trifluoromethyl),

haloaryl (e.g. chlorophenyl);

oxygen containing groups such as

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alcohols (e.g. hydroxy, hydroxyalkyl, hydroxyaryl,

20 (aryl)(hydroxy)alkyl),

ethers (e.g. alkoxy, aryloxy, alkoxyalkyl, aryloxyalkyl,

alkoxyaryl, aryloxyaryl),

aldehydes (e.g. carboxaldehyde),

ketones (e.g. alkylcarbonyl, arylcarbonyl, alkylcarbonylalkyl,

alkylcarbonylaryl, arylcarbonylalkyl, arylcarbonylaryl,

arylalkylcarbonyl, arylalkylcarbonylalkyl,

arylalkylcarbonylaryl)

acids (e.g. carboxy, carboxyalkyl, carboxyaryl),

acid derivatives such as esters

(e.g. alkoxycarbonyl, aryloxycarbonyl,

alkoxycarbonylalkyl, aryloxycarbonylalkyl,

alkoxycarbonylaryl, aryloxycarbonylaryl,

alkylcarbonyloxy, alkylcarbonyloxyalkyl),

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amides

(e.g. aminocarbonyl, mono- or di-alkylaminocarbonyl, cyclicaminocarbonyl, aminocarbonylalkyl, mono- or di-alkylaminocarbonylalkyl, arylaminocarbonyl or arylalkylaminocarbonyl, alkylcarbonylamino, arylcarbonylamino or arylalkylcarbonylamino),

carbamates

(eg. alkoxycarbonylamino, aryloxycarbonylamino, arylalkyloxycarbonylamino, aminocarbonyloxy, monoor di-alkylaminocarbonyloxy, arylaminocarbonyloxy or arylalkylaminocarbonyloxy)

and ureas

(eg. mono- or di-alkylaminocarbonylamino, arylaminocarbonylamino or

arylalkylaminocarbonylamino);

nitrogen containing groups such as

amines (e.g. amino, mono- or dialkylamino, cyclicamino, arylamino, aminoalkyl, mono- or dialkylaminoalkyl),

azides,

nitriles (e.g. cyano, cyanoalkyl),

nitro,

sulfonamides (e.g. aminosulfonyl, mono- or di-alkylaminosulfonyl, mono- or di-arylaminosulfonyl, alkyl- or aryl-sulfonyl(alkyl)amino,

alkyl- or aryl-sulfonyl(aryl)amino);

sulfur containing groups such as

thiols, thioethers, sulfoxides, and sulfones

(e.g. alkylthio, alkylsulfinyl, alkylsulfonyl, alkylsulfonyl, alkylsulfinylalkyl, alkylsulfonylalkyl, arylthio, arylsulfinyl, arylsulfonyl, arylthioalkyl, arylsulfinylalkyl, arylsulfonylalkyl)

and heterocyclic groups containing one or more, preferably one, heteroatom,

(e.g. thienyl, furanyl, pyrrolyl, imidazolyl, pyrazolyl, isothiazolyl, thiazolyl, oxazolyl, oxadiazolyl, aziridinyl, azetidinyl, thiadiazolyl, pyrrolidinyl, pyrrolinyl, imidazolidinyl, imidazolinyl, pyrazolidinyl, tetrahydrofuranyl, pyranyl, pyronyl, pyridazinyl, piperidyl, pyridyl, pyrazinyl, piperazinyl, hexahydroazepinyl, morpholinyl, thianaphthyl, benzofuranyl, isobenzofuranyl, indolyl, oxyindolyl, isoindolyl, indazolyl, indolinyl, 7azaindolyl, benzopyranyl, coumarinyl, isocoumarinyl, quinolinyl, isoquinolinyl, naphthridinyl, cinnolinyl, quinazolinyl, pyridopyridyl, benzoxazinyl, quinoxalinyl, chromenyl, chromanyl, isochromanyl, phthalazinyl and carbolinyl).

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Where any of R₁ to R₁₅ is selected from aryl or from an aryl-containing group such as aryloxy or arylthio, preferred substituent group(s) are selected from halogen, alkyl (substituted or unsubstituted; and where substituted particularly from alkoxyalkyl, hydroxyalkyl, aminoalkyl and haloalkyl), hydroxy, alkoxy, CN, NO₂, amines (including amino, mono- and dialkylamino), alkoxycarbonyl, aminocarbonyl, carboxamido, sulfonamido, alkoxycarbonylamino and aryl, and particularly from unsubstituted alkyl, substituted alkyl (including alkoxyalkyl and aminoalkyl), halogen and amines.

In one embodiment, where any of R₁ to R₁₅ is directly substituted by an alkyl substituent group, or by an alkyl-containing substituent group (such as alkoxy or alkylcarbonylamino for example), then the alkyl moiety of the substituent group directly attached to any of R₁ to R₁₅ may be further substituted by the substituent groups hereinbefore described and particularly by halogen, hydroxy, alkoxy, CN, amines (including amino, mono- and di-alkyl amino) and aryl.

In a further embodiment, where any of R₁ to R₁₅ is directly substituted by an aryl substitutent group, or by an aryl-containing substituent group (such as aryloxy or arylaminocarbonylamino for example), then the aryl moiety of the substituent group directly attached to any of R₁ to R₁₅ may be further substituted by the substituent groups hereinbefore described and particularly by

halogen, alkyl (substituted or unsubstituted; and where substituted particularly from alkoxyalkyl, hydroxyalkyl, aminoalkyl and haloalkyl), hydroxy, alkoxy, CN, NO₂, amines (including amino, mono- and di-alkylamino), alkoxycarbonyl, aminocarbonyl, carboxamido, sulfonamido, alkoxycarbonylamino and aryl. In a further embodiment, said aryl moiety is substituted by halogen, alkyl (including CF₃), hydroxy, alkoxy, CN, amines (including amino, mono- and di-alkyl amino) and NO₂. In a further embodiment, said aryl moiety is substituted by unsubstituted alkyl, substituted alkyl (particularly alkoxyalkyl and aminoalkyl), halogen and amines.

- The terms "directly substituted" and "directly attached", as used herein, mean that the substituent group is bound directly to any of R₁ to R₁₅ without any intervening divalent atoms or groups.
- In the compounds of formula (I), R₁ is selected from alkyl (including branched and unbranched alkyl, substituted and unsubstituted alkyl, and cyclic and acyclic alkyl), alkoxy, aryloxy, alkylthio, arylthio, aryl (including heteroaryl), halogen, CN, NR₇R₈ (including NH₂, mono-alkylamino and di-alkylamino), NR₆COR₇, NR₆CONR₇R₈, NR₆CO₂R₉ and NR₆SO₂R₉.
- Where R₁ is selected from alkyl, alkoxy and alkylthio, then said alkyl group or the alkyl group of the alkoxy or alkylthio is preferably selected from C₁₋₆ alkyl (including branched and unbranched alkyl, substituted and unsubstituted alkyl, and cyclic and acyclic alkyl), preferably saturated C₁₋₆ alkyl, and more preferably lower alkyl. In one embodiment, R₁ is selected from substituted alkyl, particularly haloalkyl (including CF₃) and arylalkyl (including CF₃).
 - Where R_1 is selected from NR_6COR_7 , $NR_6CONR_7R_8$, $NR_6CO_2R_9$ and $NR_6SO_2R_9$, preferably R_6 is H or alkyl, and preferably H.
- It is preferred that R₁ is selected from alkyl, alkoxy, thioalkyl, NR₇R₈ (including NH₂), NR₆COR₇, NR₆CONR₇R₈, NR₆CO₂R₉ and NR₆SO₂R₉, and preferably from NR₇R₈ (including NH₂), NR₆COR₇, NR₆CONR₇R₈, NR₆CO₂R₉ and NR₆SO₂R₉, more preferably from NR₇R₈ (including NH₂), and more preferably from NH₂.

In the compounds of formula (I), R₂ is selected from aryl (including heteroaryl) attached via an unsaturated carbon atom. R₂ may be substituted or unsubstituted. Preferably, R₂ is a 5- or 6-membered monocyclic aryl group.

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It is preferred that R₂ is not an ortho, ortho-disubstituted aryl group. Preferably, R₂ is not substituted at an ortho position relative to the point of attachment of the aryl group to the pyrimidine ring.

10 Preferably, R₂ is selected from substituted or unsubstituted heteroaryl attached via an unsaturated carbon atom.

Preferably, R₂ is a heteroaryl group which is attached to the pyrimidine ring of formula (I) such that at least one heteroatom is adjacent to the unsaturated carbon atom attached to the pyrimidine ring. Preferably, R₂ is an N, O or S-containing heteroaryl group, containing one or more heteroatom(s) selected from N, O and S.

In a preferred embodiment, R₂ is selected from furyl (including 2-furyl), thienyl (including 2-thienyl), pyridyl (including 2-pyridyl), thiazolyl (including 2- and 5- thiazolyl), pyrazolyl (including 3-pyrazolyl), triazolyl (including 4-triazolyl), pyrrolyl (including 2-pyrrolyl) and oxazolyl (including 5-oxazolyl). In a further embodiment, R₂ is selected from 2-furyl, 2-thienyl, 2-thiazolyl, 2-pyridyl, 3-pyrazolyl, 2-pyrrolyl, 4-triazolyl and 5-oxazolyl. In a further preferred embodiment, R₂ is selected from furyl, thienyl, pyridyl, thiazolyl and pyrazolyl, and particularly from 2-furyl, 2-thienyl, 2-thiazolyl, 2-pyridyl and 3-pyrazolyl. In a further embodiment, R₂ is selected from furyl, thienyl and pyridyl, preferably 2-furyl, 2-thienyl and 2-pyridyl, and more preferably from 2-furyl.

In a particularly preferred embodiment, R₂ is furyl, and preferably 2-furyl.

In the compounds of formula (I), R₃ and R₄ are independently selected from H, alkyl (including branched and unbranched alkyl, substituted and unsubstituted alkyl, and cyclic and acyclic alkyl), halogen, alkoxy, alkylthio, CN and NR₇R₈.

Where R_3 and R_4 are independently selected from alkyl, alkoxy and alkylthio, then said alkyl group or the alkyl group of the alkoxy or alkylthio is preferably selected from C_{1-6} alkyl (including branched and unbranched alkyl, substituted and unsubstituted alkyl, and cyclic and acyclic alkyl), preferably saturated C_{1-6} alkyl, and more preferably lower alkyl.

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Preferably, R₃ and R₄ are independently selected from hydrogen.

In the compounds of formula (I), R₅ is selected from H, acyclic alkyl (including branched and unbranched alkyl, and substituted and unsubstituted alkyl), COR₆, CONR₇R₈, CONR₆NR₇R₈, CO₂R₉ and SO₂R₉.

Where R_5 is selected from CONR₇R₈, R_7 and R_8 are selected from H, alkyl (including substituted alkyl such as arylalkyl (including heteroarylalkyl)) and aryl (including heteroaryl) or R_7 and R_8 may be linked to form a heterocyclic ring. In a preferred embodiment, R_7 and R_8 are selected from H, unsubstituted alkyl, arylalkyl (including heteroarylalkyl) and aryl (including heteroaryl). Said aryl groups may be substituted or unsubstituted. In a preferred embodiment one of R_7 and R_8 is hydrogen. In a further preferred embodiment, R_7 is H and R_8 is selected from arylalkyl (including heteroarylalkyl), preferably arylmethyl (including heteroarylmethyl).

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Where R_5 is selected from acyclic alkyl, it is preferably C_{1-6} acyclic alkyl (including alkenyl and alkynyl). In one embodiment, R_5 is selected from C_{1-6} saturated acyclic alkyl, preferably lower alkyl.

- In one embodiment, R₅ is selected from substituted acyclic alkyl (including saturated acyclic alkyl and alkenyl). Preferred substituents are aryl (including heteroaryl), cycloalkyl, non-aromatic heterocyclyl, CO₂R₆, CONR₇R₈, CONR₆NR₇R₈ and C(=NR₆)NR₇R₈, preferably aryl (including heteroaryl) and CONR₇R₈, and more preferably aryl (including heteroaryl).
- Where R₅ is selected from acyclic alkyl substituted by aryl (including heteroaryl), the aryl (including heteroaryl) group may be substituted or unsubstituted, preferably substituted. Preferred aryl groups are discussed below with regard to the group R₁₂. Preferably, the aryl-substituted acyclic alkyl is an aryl-substituted methyl group.

In one embodiment, R_5 is selected from $(CR_{10}R_{11})_nR_{12}$ wherein n is 1 to 6 (preferably n is 1), R_{10} and R_{11} are independently selected from H, alkyl and aryl, and R_{12} is selected from aryl (including heteroaryl), cycloalkyl, non-aromatic heterocyclic, CO_2R_6 , $CONR_7R_8$, $CONR_6NR_7R_8$ and $C(=NR_6)NR_7R_8$. Preferably, R_{12} is selected from aryl (including heteroaryl) and $CONR_7R_8$, and more preferably from aryl (including heteroaryl). Preferably R_{10} and R_{11} are independently selected from H and alkyl, more preferably H.

Where R_{12} is selected from CONR₇R₈, R_7 and R_8 are selected from H, alkyl (including substituted alkyl such as arylalkyl (including heteroarylalkyl)) and aryl (including heteroaryl) or R_7 and R_8 may be linked to form a heterocyclic ring. In one embodiment, R_7 and R_8 are selected from H, unsubstituted alkyl, arylalkyl (including heteroarylalkyl) and aryl (including heteroaryl). Said aryl groups may be substituted or unsubstituted. In a preferred embodiment one of R_7 and R_8 is hydrogen.

- Where R_{12} is selected from aryl (including heteroaryl), the aryl group may be unsubstituted or substituted, and is preferably substituted. In a preferred embodiment, R₁₂ is selected from mono-, di- or tri-substituted aryl (including heteroaryl) groups. Where R₁₂ is heteroaryl, R₁₂ is preferably selected from mono or bicyclic heteroaryl groups, more preferably from pyridyl ... (including 2-pyridyl, 3-pyridyl and 4-pyridyl, preferably 2-pyridyl), indolyl (including 2indolyl, 3-indolyl, 4-indolyl, 5-indolyl, 6-indolyl and 7-indolyl), furyl (including 2-furyl and 3furyl, preferably 2-furyl), thienyl (including 2-thienyl and 3-thienyl, preferably 2-thienyl), isoindolyl, indolinyl, isoxazolyl, oxazolyl, thiazolyl, pyrazinyl, pyrimidinyl, quinolinyl, benzoxadiazolyl, benzothiadiazolyl, benzotriazolyl, indazolyl, benzodioxolyl and dihydrobenzofuranyl, more preferably from pyridyl (preferably 2-pyridyl), indolyl, furyl (preferably 2-furyl) and thienyl (preferably 2-thienyl), and most preferably from pyridyl (preferably 2-pyridyl), furyl (preferably 2-furyl) and thienyl (preferably 2-thienyl). Preferably, R_{12} is selected from phenyl, thienyl, furyl and pyridyl, more preferably from phenyl, 2-thienyl, 2-furyl and 2-pyridyl.
- In one embodiment, R_{12} is selected from mono-, di- or tri-substituted aryl (including heteroaryl) groups represented by the formula $Ar(R_{13})_a(R_{14})_b(R_{15})_c$ wherein Ar is an aryl (including heteroaryl) group, preferably selected from the preferred aryl groups described

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above for R_{12} ; wherein R_{13} , R_{14} and R_{15} are substituent group(s), the same or different; and wherein a, b and c are 0 or 1 such that a+b+c ≥ 1 .

The substituent groups R_{13} , R_{14} and R_{15} may be selected from any of the substituent groups described herein above.

In a preferred embodiment, R₁₃, R₁₄ and R₁₅ are selected from NR₇R₈ (including NH₂, and NHR₇) alkyl (substituted or unsubstituted; preferably C₁₋₆ acyclic alkyl), alkoxy (including fluoroalkoxy), halogen (including F, Cl, Br and I), NO₂, CN, hydroxy, NHOH, CHO, CONR₇R₈, CO₂R₇, NR₆COR₇ (preferably NHCOR₇), NR₆CO₂R₉ (preferably NHCO₂R₉), NR₆SO₂R₉ (preferably NHSO₂R₉), OCO₂R₉ and aryl (including heteroaryl).

In a more preferred embodiment, R₁₃, R₁₄ and R₁₅ are selected from NR₇R₈ (including NH₂ and NHR₇), alkyl (substituted or unsubstituted; and preferably C₁₋₆ acyclic saturated alkyl) and halogen (preferably F or Cl, particularly F).

In a particularly preferred embodiment, R_{13} , R_{14} and R_{15} are selected from NR_7R_8 (including NH_2 and NHR_7 , preferably NH_2) and alkyl (substituted or unsubstituted; preferably C_{1-6} acyclic saturated alkyl).

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Where R₁₃, R₁₄ and R₁₅ are selected from substituted alkyl, said alkyl is preferably selected from alkoxyalkyl, hydroxyalkyl, aminoalkyl (including NH₂-alkyl, mono-alkylaminoalkyl and di-alkylaminoalkyl), haloalkyl (particularly fluoroalkyl (including CF₃)), cyanoalkyl, alkylthioalkyl, alkylcarboxyaminoalkyl, alkoxycarbonylaminoalkyl and alkylsulfonylamino, more preferably from alkoxyalkyl, hydroxyalkyl, aminoalkyl and haloalkyl (particularly fluoroalkyl (including CF₃)) and most preferably from alkoxyalkyl and aminoalkyl.

In one embodiment, the substituent groups R₁₃, R₁₄ and R₁₅ are selected from halogen, alkyl (including CF₃), hydroxy, alkoxy, alkylthio, CN, amines (including amino, mono- and di-alkyl amino) and NO₂.

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In a preferred embodiment, R₅ is selected from H and substituted acyclic alkyl, preferably wherein said acyclic alkyl is substituted by aryl (including heteroaryl) or CONR₇R₈.

In the compounds of formula (I), R_6 , R_7 , R_8 , R_{10} and R_{11} are independently selected from H, alkyl (including branched and unbranched alkyl, substituted and unsubstituted alkyl, cyclic and acyclic alkyl) and aryl (including heteroaryl) or where R_7 and R_8 are in any NR_7R_8 group R_7 and R_8 may be linked to form a heterocyclic group, or where R_6 , R_7 and R_8 are in a (CONR₆NR₇R₈) group, R_6 and R_7 may be linked to form a heterocyclic group.

In the compounds of formula (I), R₉ is selected from alkyl (including branched and unbranched alkyl, substituted and unsubstituted alkyl, cyclic and acyclic alkyl) and aryl (including heteroaryl).

Where R_6 to R_{11} are independently selected from alkyl, preferably R_6 to R_{11} are selected from C_{1-6} alkyl, preferably C_{1-6} saturated alkyl and more preferably from lower alkyl.

Where R₇ and R₈, or R₆ and R₇, are linked to form a heterocyclic ring said heterocyclic ring may be saturated, partially unsaturated or aromatic, and is preferably saturated. Said heterocyclic ring is preferably a 5, 6 or 7-membered ring, preferably a 5 or 6-membered ring, and may contain one or more further heteroatom(s) preferably selected from N, O and S.

In a preferred embodiment, R₁ is NH₂, R₂ is 2-furyl, R₃ and R₄ are H and R₅ is arylmethyl (including heteroarylmethyl).

In a particularly preferred embodiment of the invention, the compound of formula (I) is 7-(2-fluorobenzyl)-4-(2-fluorobenzyl)-7H-pyrrolo[2,3-d]pyrimidine-2-amine.

Where chiral the compounds of formula (I) may be in the form of a racemic mixture of pairs of enantiomers or in enantiomerically pure form.

According to a further aspect of the present invention there is provided a method of treating or preventing a disorder in which the blocking of purine receptors, particularly adenosine receptors and more particularly adenosine A_{2A} receptors, may be beneficial, the method

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comprising administration to a subject in need of such treatment an effective dose of a compound of formula (I) or a pharmaceutically acceptable salt or prodrug thereof.

The disorder may be caused by the hyperfunctioning of the purine receptors.

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The present invention may be employed in respect of a human or animal subject, more preferably a mammal, more preferably a human subject.

The disorders of particular interest are those in which the blocking of purine receptors, partiucularly adenosine receptors and more particularly adenosine A_{2A} receptors, may be beneficial. These may include movement disorders such as Parkinson's disease, druginduced Parkinsonism, post-encephalitic Parkinsonism, Parkinsonism induced by poisoning (for example MPTP, manganese, carbon monoxide) and post-traumatic Parkinson's disease (punch-drunk syndrome).

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Other movement disorders in which the blocking of purine receptors, may be of benefit include progressive supernuclear palsy, Huntingtons disease, multiple system atrophy, corticobasal degeneration, Wilsons disease, Hallerrorden-Spatz disease, progressive pallidal atrophy, Dopa-responsive dystonia-Parkinsonism, spasticity or other disorders of the basal ganglia which result in abnormal movement or posture. The present invention may also be effective in treating Parkinson's with on-off phenomena; Parkinson's with freezing (end of dose deterioration); and Parkinson's with prominent dyskinesias.

The compounds of formula (I) may be used or administered in combination with one or more additional drugs useful in the treatment of movement disorders, such as L-DOPA or a dopamine agonist, the components being in the same formulation or in separate formulations for administration simultaneously or sequentially.

Other disorders in which the blocking of purine receptors, particularly adenosine receptors and more particularly adenosine A_{2A} receptors may be beneficial include acute and chronic pain; for example neuropathic pain, cancer pain, trigeminal neuralgia, migraine and other conditions associated with cephalic pain, primary and secondary hyperalgesia, inflammatory pain, nociceptive pain, tabes dorsalis, phantom limb pain, spinal cord injury

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pain, central pain, post-herpetic pain and HIV pain; affective disorders including mood disorders such as bipolar disorder, seasonal affective disorder, depression, manic depression, atypical depression and monodepressive disease; central and peripheral nervous system degenerative disorders including corticobasal degeneration, demyelinating disease (multiple sclerosis, disseminated sclerosis), Freidrich's ataxia, motoneurone disease (amyotrophic lateral sclerosis, progressive bulbar atrophy), multiple system atrophy, myelopathy, radiculopathy, peripheral neuropathy (diabetic neuropathy, tabes dorsalis, drug-induced neuropathy, vitamin deficiency), systemic lupus erythamatosis, granulomatous disease, olivo-ponto-cerebellar atrophy, progressive pallidal atrophy, progressive supranuclear palsy, spasticity; schizophrenia and related pyshoses; cognitive disorders including dementia, Alzheimers Disease, Frontotemporal dementia, multi-infarct dementia, AIDS dementia, dementia associated with Huntingtons Disease, Lewy body dementia, senile dementia, age-related memory impairment, cognitive impairment associated with dementia, Korsakoff syndrome, dementia pugilans; attention disorders such as attention-deficit hyperactivity disorder (ADHD), attention deficit disorder, minimal brain dysfunction, brain-injured child syndrome, hyperkinetic reaction childhood, and hyperactive child syndrome; central nervous system injury including traumatic brain injury, neurosurgery (surgical trauma), neuroprotection for head injury, raised intracranial pressure, cerebral oedema, hydrocephalus, spinal cord injury; cerebral ischaemia including transient ischaemic attack, stroke (thrombotic stroke, ischaemic stroke, embolic stroke, haemorrhagic stroke, lacunar stroke) subarachnoid haemorrhage, cerebral vasospasm, neuroprotection for stroke, peri-natal asphyxia, drowning, cardiac arrest, subdural haematoma; myocardial ischaemia; muscle ischaemia; sleep disorders such as hypersomnia and narcolepsy; eye disorders such as retinal ischaemia-reperfusion injury and diabetic neuropathy; cardiovascular disorders such as claudication and hypotension; and diabetes and its complications.

According to a further aspect of the present invention there is provided use of a compound of formula (I) or a pharmaceutically acceptable salt or prodrug thereof in the manufacture of a medicament for the treatment or prevention of movement disorders in a subject.

According to a further aspect of the invention there is provided a method of treating or preventing movement disorders comprising administration to a subject in need of such

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treatment an effective dose of a compound of formula (I) or a pharmaceutically acceptable salt or prodrug thereof.

According to a further aspect of the invention there is provided use of a compound of formula (I) or a pharmaceutically acceptable salt or prodrug thereof in the manufacture of a medicament for neuroprotection in a subject.

According to a further aspect of the invention there is provided a method of neuroprotection comprising administration to a subject in need of such treatment an effective dose of a compound of formula (I) or a pharmaceutically acceptable salt or prodrug thereof.

The medicament for or method of neuroprotection may be of use in the treatment of subjects who are suffering from or at risk from a neurodegenerative disorder, such as a movement disorder.

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According to a further aspect of the invention, there is provided for use in therapy a compound of formula (I), or a pharmaceutically acceptable salt or prodrug thereof.

According to a further aspect of the invention, there is provided a compound of formula (I), or a pharmaceutically acceptable salt or prodrug thereof, per se, other than:

- (i) compounds wherein R₂ is selected from pyrrolopyrimidines; and
- (ii) compounds wherein R_1 is selected from methyl, phenyl and (methyl)phenyl, R_2 is selected from phenyl (substituted or unsubstituted) and R_5 is selected from methyl and methanesulfonyl, preferably compounds wherein R_1 is selected from methyl and phenyl (substituted or unsubstituted), R_2 is selected from phenyl (substituted or unsubstituted) and R_5 is selected from methyl and methanesulfonyl, and more preferably wherein R_1 is selected from methyl and phenyl (substituted) and R_2 is selected from phenyl (substituted) and R_2 is selected from phenyl (substituted).
- According to a further aspect of the invention, there is provided a compound of formula (I), or a pharmaceutically acceptable salt or prodrug thereof, per se, wherein R_2 is selected from heteroaryl, other than compounds wherein R_2 is selected from pyrrolopyrimidines.

In a preferred embodiment, there is provided a compound of formula (I), or a pharmaceutically acceptable salt or prodrug thereof, *per se*, wherein R₂ is selected from 5-or 6- membered monocyclic heteroaryl groups.

According to a further aspect of the invention, there is provided a method of preparing the novel compounds of the present invention. Compounds of formula (I) may be prepared according to conventional synthetic methods, such as set out in Reaction Scheme 1.

Reaction Scheme 1

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Compounds of formula (1) where R₁ is alkoxy, aryloxy, alkylthio, arylthio, CN or NR₇R₈ may be prepared from compounds of formula (2) by standard methods such as nucleophilic displacement using an appropriate nucleophilic reagent such as an alcohol, thiol, cyanide or amine (HNR₇R₈) in the presence of a suitable base if required.

Compounds of formula (1) where R₁ is NR₆COR₇, NR₆CO₂R₉ or NR₆SO₂R₉, wherein R₆ is H, alkyl or aryl, may be prepared from compounds of formula (1) where R₁ is NR₇R₈, wherein R₇ is H and R₈ is H, alkyl or aryl, by standard methods such as treatment with an appropriate acid chloride (R₇COCl), chloroformate (ClCO₂R₉) or sulphonyl chloride (R₉SO₂Cl) in the presence of a suitable base such as triethylamine.

Compounds of formula (1) where R₁ is NR₆CONR₇R₈ wherein R₆ is H, alkyl or aryl, may be prepared from compounds of formula (1) where R₁ is NR₇R₈, wherein R₇ is H and R₈ is

H, alkyl or aryl, by standard methods such as treatment with an appropriate isocyanate (R₇NCO or R₈NCO) or carbamoyl chloride (R₇R₈NCOCl) in the presence of a suitable base such as triethylamine if required.

Compounds of formula (1) where R₁ is NH₂ may be prepared from compounds of formula (1) where R₁ is NR₇R₈, wherein R₇ is H and R₈ is a suitable protecting group such as a 3,4-dimethoxybenzyl group, by standard methods such as treatment with TFA.

Compounds of formula (2) where R_5 is alkyl (including arylalkyl, heteroarylalkyl and $(CR_{10}R_{11})_nR_{12}$) may be prepared from a compound of formula (3) by standard methods such as reaction with an appropriate alkyl halide, or substituted alkyl halide in the presence of a suitable base such as sodium hydride.

Compounds of formula (2) where R₅ is $(CR_{10}R_{11})_nR_{12}$ and R₁₂ is $CONR_7R_8$ or $CONR_6NR_7R_8$ may be prepared from compounds of formula (2) where R₅ is $(CR_{10}R_{11})_nR_{12}$ and R₁₂ is CO_2R_6 by standard methods such as direct reaction with an appropriate amine or hydrazine or by initial hydrolysis of the ester group CO_2R_6 to a carboxylic acid followed by reaction with an appropriate amine or hydrazine in the presence of a standard coupling reagent such as DCC.

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Compounds of formula (2) where R_5 is $(CR_{10}R_{11})_nR_{12}$, wherein R_{12} is $C(=NR_6)NR_7R_8$ may be prepared from compounds of formula (2) where R_5 is $(CR_{10}R_{11})_nR_{12}$ wherein R_{12} is CN by standard methods such as treatment with an appropriate amine in the presence of trimethyl aluminium.

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Compounds of formula (2) where R_5 is $(CR_{10}R_{11})_nR_{12}$, wherein R_{12} is CO_2R_6 or CN may be prepared from compounds of formula (3) by standard methods such as treatment with an appropriate substituted alkyl halide in the presence of a suitable base such as sodium hydride.

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Compounds of formula (2) where R₅ is CONR₇R₈ or CONR₆NR₇R₈ may be prepared from compounds of formula (3) by standard methods such as treatment with an appropriate isocyanate (R₇NCO or R₈NCO) or carbamoyl chloride (R₇R₈NCOCl or R₇R₈NR₆NCOCl).

Compounds of formula (2) where R₅ is COR₆, CO₂R₉ or SO₂R₉ may be prepared from compounds of formula (3) by standard methods such as treatment with an appropriate acid chloride (R₆COCl), chloroformate (ClCO₂R₉) or sulphonyl chloride (R₉SO₂Cl) in the presence of a suitable base such as triethylamine.

Compounds of formula (3) are prepared from compounds of formula (4) by standard methods such as aryl or heteroaryl coupling reactions. Suitable aryl or heteroaryl coupling reactions would include reaction with an appropriate aryl or heteroaryl trialkylstannane derivative, an aryl or heteroarylboronic acid or boronic ester derivative, or an aryl or heteroarylzinc halide derivative in the presence of a suitable catalyst such as a palladium complex. Compounds of formula (4) are either known in the literature or are prepared by methods which are analogous to those described in the literature.

In certain cases it may be advantageous to prepare a compound of formula (2) where R₅ is selected to perform the function of a protecting group, for example a suitable protecting group would be a benzyl group or substituted benzyl group such as a 3,4-dimethoxybenzyl group or a trimethylsilylethoxymethyl group. Compounds of this nature may prepared as described above and converted to compounds of formula (1), where R₅ is a protecting group, as described above. The protecting group R₅ may be then be removed by standard methods such as treatment with, for example, TFA or tetra-*n*-butylammonium fluoride to give a compound of formula (1) where R₅ is H. Compounds of formula (1) where R₅ is H may then be used to prepare other compounds of formula (1), where R₅ is as previously defined, by the methods described above.

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In certain cases it may be advantageous to prepare compounds of formula (1) where R_5 is alkyl (including arylalkyl, heteroarylalkyl and $(CR_{10}R_{11})_nR_{12}$ from compounds of formula (1) where R_5 is H by the methods described above.

Other compounds of formula (1) may be prepared by standard methods such as those illustrated in Reaction Scheme 2.

Reaction Scheme 2

Compounds of formula (1) where R_1 is alkyl or aryl may be prepared from compounds of formula (5) where R_1 is alkyl or aryl by standard methods such as aryl or heteroaryl coupling reactions as described above. Compounds of formula (5) where R_1 is alkyl or aryl are either known in the literature or may be prepared by methods analogous to those described in the literature.

Compounds of formula (1) where R₃ and R₄ are H, alkyl, halogen, alkoxy, alkylthio, CN or NR₇R₈ may be prepared by the methods described above. Suitable intermediates such as compounds of formula (4) and (5) where R₃ and R₄ are as described above, are either known in the literature or are prepared by methods which are analogous to those described in the literature.

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According to a further aspect of the invention, there is provided a pharmaceutical composition comprising a compound of formula (I) in combination with a pharmaceutically acceptable carrier or excipient and a method of making such a composition comprising combining a compound of formula (I) with a pharmaceutically acceptable carrier or excipient.

The pharmaceutical compositions employed in the present invention comprise a compound of formula (I), or pharmaceutically acceptable salts or prodrugs thereof, and may also contain a pharmaceutically acceptable carrier and optionally other therapeutic ingredients known to those skilled in the art. The term, "pharmaceutically acceptable salts", refers to salts prepared from pharmaceutically acceptable non-toxic acids including inorganic acids and organic acids.

Where the compounds of formula (I) are basic, salts may be prepared from 30 pharmaceutically acceptable non-toxic acids including inorganic and organic acids. Such

acids include acetic, benzenesulfonic, benzoic, camphorsulfonic, citric, ethenesulfonic, fumaric, gluconic, glutamic, hippuric, hydrobromic, hydrochloric, isethionic, lactic, maleic, malic, mandelic, methanesulfonic, mucic, nitric, pamoic, pantothenic, phosphoric, succinic, sulfuric, tartaric, oxalic, p-toluenesulfonic and the like. Particularly preferred are hydrochloric, hydrobromic, phosphoric, and sulfuric acids, and most particularly preferred is the hydrochloride salt.

Any suitable route of administration may be employed for providing the patient with an effective dosage of a compound of formula (I). For example, oral, rectal, parenteral (intravenous, intramuscular), transdermal, subcutaneous, and the like may be employed. Dosage forms include tablets, troches, dispersions, suspensions, solutions, capsules, patches, and the like. The most suitable route in any given case will depend on the severity of the condition being treated. The most preferred route of administration of the present invention is the oral route. The compositions may be conveniently presented in unit dosage form and prepared by any of the methods well known in the art of pharmacy.

In practical use, the compounds of formula (I) can be combined as the active ingredient in intimate admixture with a pharmaceutical carrier according to conventional pharmaceutical compounding techniques. The carrier may take a wide variety of forms depending on the form of preparation desired for administration, e.g. oral or parenteral (e.g. intravenous). In preparing the compositions for oral dosage form, any of the usual pharmaceutical media may be employed as carriers, such as, for example, water, glycols, oils, alcohols, flavouring agents, preservatives, colouring agents, and the like in the case of oral liquid preparations (such as suspensions, solutions and elixirs) or aerosols; or carriers such as starches, sugars, micro-crystalline cellulose, diluents, granulating agents, lubricants, binders, disintegrating agents, and the like may be used in the case of oral solid preparations such as, for example, powders, capsules, and tablets, with the solid oral preparation being preferred over the liquid preparations. The most preferred solid oral preparation is tablets.

30 Because of their ease of administration, tablets and capsules represent the most advantageous oral dosage unit form in which case solid pharmaceutical carriers are employed. If desired, tablets may be coated by standard aqueous or non-aqueous techniques.

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In addition to the common dosage forms set out above, the compounds of formula (I) may also be administered by controlled release means and/or delivery devices such as those described in United States Patent Nos.: 3,845,770; 3,916,899; 3,536,809; 3,598,123; 3,630,200; 4,008,719; 4,687,660; and 4,769,027, the disclosures of which are hereby incorporated by reference.

Pharmaceutical compositions employed in the present invention suitable for oral administration may be presented as discrete units such as capsules, cachets, or tablets, or aerosol sprays each containing a predetermined amount of the active ingredient as a powder or granules, a solution or a suspension in an aqueous liquid, an oil-in-water emulsion, or a water-in-oil liquid emulsion. Such compositions may be prepared by any of the methods of pharmacy, but all methods include the step of bringing the active ingredient into association with the carrier which constitutes one or more necessary ingredients. In general, the compositions are prepared by uniformly and intimately admixing the active ingredient with liquid carriers or finely divided solid carriers or both, and then, if necessary, shaping the product into the desired presentation.

For example, a tablet may be prepared by compression or moulding, optionally with one or more accessory ingredients. Compressed tablets may be prepared by compressing in a suitable machine the active ingredient in a free-flowing form such as powder or granules, optionally mixed with a binder, a lubricant, an inert diluent, and/or a surface active or dispersing agent. Moulded tablets may be made by moulding in a suitable machine a mixture of the powdered compound moistened with an inert liquid diluent.

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The invention is further defined by reference to the following examples. It will be apparent to those skilled in the art that many modifications, both to materials and methods, may be practised without departing from the purpose and interest of this invention.

EXAMPLES 30

Synthetic Examples

The invention is illustrated with reference to the following Examples, as set out in Table 1.

Table 1

Example	Structure	Compound Name
1	N N NMe ₂	N,N-dimethyl-4-(2-furyl)-1H-pyrrolo[2,3-d]pyrimidine-2-amine
2	NH CI	2-chloro-4-(2-furyl)-1H-pyrrolo[2,3-d]pyrimidine
3	N N NMe ₂	7-benzyl-N,N-dimethyl-4-(2-furyl)-7H-pyrrolo[2,3-d]pyrimidine-2-amine
4	N N CI	7-benzyl-2-chloro-4-(2-furyl)-7H-pyrrolo[2,3-d]pyrimidine
5	N N NMe ₂	7-benzoyl-N,N-dimethyl-4-(2-furyl)-7H-pyrrolo[2,3-d]pyrimidine-2-amine
6	HON OH	(2R)-2-(2-hydroxymethylpyrrolidin-1-yl)-4- (2-furyl)-1H-pyrrolo[2,3-d]pyrimidine
7	N N OMe OMe	N-(3,4-dimethoxybenzyl)-4-(2-furyl)-1H-pyrrolo[2,3-d]pyrimidine-2-amine
8	N NH ₂	4-(2-furyl)-1H-pyrrolo[2,3-d]pyrimidine-2-amine

9	HAN	N-benzyl-N'-(4-(2-furyl)-1H-pyrrolo[2,3-d]pyrimidin-2-yl)urea
10	F CI	2-chloro-7-(2-fluorobenzyl)-4-(2-furyl)-7H-pyrrolo[2,3-d]pyrimidine
11	N N NMe ₂	N,N-dimethyl-7-(2-fluorobenzyl)-4-(2-furyl)-7H-pyrrolo[2,3-d]pyrimidine-2-amine
12	OMe OMe	N-(3,4-dimethoxybenzyl)-7-(2-fluorobenzyl)-4-(2-furyl)-7H-pyrrolo[2,3-d]pyrimidine-2-amine
13	N N NH ₂	7-(2-fluorobenzyl)-4-(2-furyl)-7H-pyrrolo[2,3-d]pyrimidine-2-amine

The syntheses of the above Examples is described with reference to the following general synthetic Methods. The analytical data for the Examples, together with the Method of synthesis used is given in Table 2.

5

Synthetic Methods

Method A

2-Chloro-4-(2-furyl)-1H-pyrrolo[2,3-d]pyrimidine (Example 2)

A solution of 2,4-dichloro-1H-pyrrolo[2,3-d]pyrimidine (4.1 g, 21.8 mmol) in DMF (20 mL) was treated with PdCl₂(PPh₃)₂ (772 mg, 0.17 mmol) and 2-(tributylstannyl)-furan (6.9 mL, 21.8 mmol), stirred at room temperature for 16 h, diluted with diethyl ether and filtered to give the *title compound* (2.94 g, 61 %) as a pale orange solid.

15 Method B

2-chloro-7-(2-fluorobenzyl)-4-(2-furyl)-7H-pyrrolo[2,3-d]pyrimidine (Example 10)

A solution of 2-chloro-4-(2-furyl)-1H-pyrrolo[2,3-d]pyrimidine (219 mg, 1 mmol) in DMF (2 mL) at 0 °C was treated with NaH (40 mg, 60 %, 1 mmol), stirred for 20 min, treated with 2-fluorobenzyl bromide (120 μL, 1 mmol), stirred at room temperature for 1 h, quenched with water, extracted with EtOAc, dried (MgSO₄), concentrated *in vacuo* and purified by chromatography (EtOAc : Heptane, 1:4) to give the *title compound* (250 mg, 76 %) as a cream solid.

Method C

N-(3,4-dimethoxybenzyl)-7-(2-fluorobenzyl)-4-(2-furyl)-7H-pyrrolo[2,3-d]pyrimidine-

10 **2-amine (Example 12)**

A solution of 2-chloro-7-(2-fluorobenzyl)-4-(2-furyl)-7H-pyrrolo[2,3-d]pyrimidine (254 mg, 0.78 mmol) in N-methylpyrrolidone (2 mL) was treated with veratrylamine (0.25 mL, 1.66 mmol), heated to 100 °C for 16 h and purified by chromatography (Heptane : EtOAc, 4:1) to give the title compound (316 mg, 88 %) as a cream solid.

15

Method D

7-(2-fluorobenzyl)-4-(2-furyl)-7H-pyrrolo[2,3-d]pyrimidine-2-amine (Example 13)

A solution of N-(3,4-dimethoxybenzyl)-7-(2-fluorobenzyl)-4-(2-furyl)-7H-pyrrolo[2,3-d]pyrimidine-2-amine (110 mg, 0.24 mmol) in TFA (1 mL) was heated to 50 °C for 3 h, concentrated *in vacuo*, treated with saturated NaHCO₃, extracted with EtOAc, dried (MgSO₄), concentrated *in vacuo*, purified by chromatography (EtOAc: Heptane, 1:4) and recrystallised (EtOAc) to give the *title compound* (38 mg, 51 %) as a cream solid.

Method E

7-Benzoyl-N,N-dimethyl-4-(2-furyl)-7H-pyrrolo[2,3-d]pyrimidine-2-amine (Example 5)

A solution of N,N-dimethyl-4-(2-furyl)-1H-pyrrolo[2,3-d]pyrimidine-2-amine (181 mg, 0.8 mmol) in THF (3 mL) was treated with Et₃N, (111 μL, 0.8 mmol), benzoyl chloride (93 μL, 0.8 mmol) and a catalytic amount of DMAP, stirred at room temperature for 16 h, quenched with water, extracted with EtOAc, dried (MgSO₄), concentrated *in vacuo* and purified by chromatography (EtOAc: Heptane, 1:9 then Heptane: DCM, 2:1) to give the title compound (76 mg) as a yellow solid.

Method F

N-benzyl-N'-(4-(2-furyl)-1H-pyrrolo[2,3-d]pyrimidin-2-yl)urea (Example 9)

A solution of 4-(2-furyl)-1H-pyrrolo[2,3-d]pyrimidine-2-amine (192 mg, 0.96 mmol) in THF (3 mL) was treated with benzyl isocyanate (118 µL, 0.96 mmol), stirred at room temperature for 16 h and the resulting solid filtered and washed with EtOAc to give the *title compound* (61 mg, 19 %) as a yellow solid.

Method G

2,4-Dichloro-7-(2-trimethylsilylethoxymethyl)-7H-pyrrolo[2,3-d]pyrimidine

A solution of 2,4-dichloro-1H-pyrrolo[2,3-d]pyrimidine (542 mg, 2.58 mmol) in MeCN (20 mL) was treated with NaH (113 mg, 2.84 mmol), stirred for 1 h, treated with trimethylsilylethyl chloride (502 μL, 2.84 mmol), stirred at room temperature for 3 h, poured into water, extracted with EtOAc, dried (MgSO₄) and concentrated *in vacuo* to give the title compound (913 mg, 104 %) which was used crude in the next reaction.

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Method H

N,N-Dimethyl-4-(2-furyl)-1H-pyrrolo[2,3-d]pyrimidine-2-amine (Example 1)

A solution of N,N-dimethyl-4-(2-furyl)-7-(2-trimethylsilylethoxymethyl)-7H-pyrrolo[2,3-d]pyrimidine-2-amine (370 mg, 0.97 mmol) in THF (3 mL) was treated with

- tetrabutylammonium fluoride (1 ml, 1-M in THF, 1 mmol),refluxed for 36 h, poured into water, extracted with EtOAc, dried (MgSO₄), concentrated in vacuo and purified by chromatography (EtOAc: Heptane, 1:9, alumina) to give the *title compound* (47 mg, 21%) as a yellow solid.
- 25 2-Chloro-4-(2-furyl)-7-(2-trimethylsilylethoxymethyl)-7H-pyrrolo[2,3-d]pyrimidine
 This was prepared from 2,4-dichloro-7-(2-trimethylsilylethoxymethyl)-7H-pyrrolo[2,3-d]pyrimidine by method A and the *title compound* (207 mg, 21 %) isolated as a cream solid.
- N,N-Dimethyl-4-(2-furyl)-7-(2-trimethylsilylethoxymethyl)-7H-pyrrolo[2,3-d]pyrimidine-2-amine

This was prepared from 2-chloro-4-(2-furyl)-7-(2-trimethylsilylethoxymethyl)-7H-pyrrolo[2,3-d]pyrimidine by method C and the *title compound* (409 mg, 77 %) isolated as a white solid.

Table 2

Example	Method	Yield(%)	Physical Data
			mp 196.9– 197.1 °C; NMR δ_H (400 MHz, CDCl ₃) 3.27 (6H, s), 6.57 –
1	H	21	6.60 (1H, m), 6.83 – 6.86 (1H, m), 6.93 – 6.97 (1H, m), 7.31 (1H, d, J 4.0
			Hz), 7.65 – 7.68 (1H, m), 8.84 (1H, s)
		61	mp 265.8 – 266.3 °C; IR v_{max} (Nujol)/cm ⁻¹ 3199, 3108, 2924, 2854,
2	A		1593, 1530 and 1462; NMR $\delta_{\rm H}$ (400 MHz, DMSO) 6.79 – 6.82 (1H, m),
			6.98 – 6.99 (1H. m), 7.50 (1H, d, J 3.5 Hz), 7.67 (1H, t, J 3.5 Hz), 8.08
			(1H, d. J 1.0 Hz), 12.4 (1H, s)
			mp 91.7 – 92.3 °C; IR v_{max} (Nujol)/cm ⁻¹ 3020, 2924, 2854, 1613, 1582,
3	В	•	1542, 1452, 1395 and 733; NMR δ_{H} (400 MHz, CDCl ₃) 3.27 (6H, s),
			5.31 (2H, s), 6.55 – 6.57 (1H, m), 6.80 (1H, d, J 3.5 Hz), 6.81 (1H, d, J
			4.0 Hz), 7.23 – 7.34 (6H, m), 7.63 – 7.65 (1H, m);
			IR v _{max} (Nujol)/cm ⁻¹ 3116, 3064, 3032, 2931, 1600, 1565, 1509, 1328,
4	В		1256, 1151 and 920; NMR δ_{H} (400 MHz, CDCl ₃) 5.43 (2H, s), 6.61 –
	ם		6.65 (1H, m), 7.01 (1H, d, J 3.5 Hz), 7.15 (1H, d, J 3.5 Hz), 7.21 – 7.37
			(5H, m), 7.47 (1H, dd, J 3.5, 1.0 Hz), 7.69 – 7.71 (1H, m)
			mp 108.3 – 109.1 °C; NMR $\delta_{\rm H}$ (400 MHz, CDCl ₃) 2.89 (6H, s), 6.57 –
			6.60 (1H, m), 7.05 (1H, d, J 4.0 Hz), 7.28 – 7.29 (1H, m), 7.43 – 7.48
5	E		(2H, m), 7.53 – 7.59 (2H, m), 7.66 – 7.67 (1H, m), 7.75 – 7.79 (2H, m);
			Anal. Calcd for $C_{19}H_{16}N_4O_2 + 0.3 H_2O$: C, 67.56; H, 4.95, N, 16.59.
			Found: C, 67.48; H, 4.72; N, 16.45; .M/Z 333 (M+H) ⁺ .
·	С	64	mp 174.8 – 175.6 °C; IR v_{max} (Nujol)/cm ⁻¹ 3094, 2752, 1611, 1560, 1518
			and 1467; NMR δ_H (400 MHz, DMSO) 1.81 – 2.04 (2H, m), 3.26 – 3.36
6			(2H, m), 3.56 – 3.47 (1H, m), 3.47 – 3.62 (2H, m), 3.69 – 3.76 (1H, m),
			4.11 – 4.24 (1H, m), 5.01 (1H, s), 6.66 – 6.69 (1H, m), 6.71 – 6.74 (1H,
			m), 7.11 (1H, dd, J 3.5, 2.0 Hz), 7.30 (1H, dd, J 3.5, 1.0 Hz), 7.96 – 7.97
			(1H, m), 11.41 (1H, s).

		<u></u>	
7			mp 154.8 – 154.9 °C; IR v_{max} (Nujol)/cm ⁻¹ 3459, 3416, 3142, 1623,
			1593, 1482, 1467, 1266 and 742; NMR $\delta_{\rm H}$ (400 MHz, DMSO) 3.69 (3H,
	C	82	s), 3.71 (3H, s), 4.47 (2H, d, J 6.0 Hz), 6.65 – 6.67 (1H, m), 6.71 – 6.72
			(1H, m), 6.83 – 6.91 (2H, m), 7.03 – 7.09 (3H, m), 7.27 (1H, d, J 2.5 Hz),
			$7.94 - 7.96$ (1H, m), 11.27 (1H, s); Anal. Calcd for $C_{19}H_{18}N_4O_3 + 2 H_2O$:
			C, 59.06; H, 5.749, N, 14.50. Found: C, 59.07; H, 5.74; N, 14.19.
			mp 213.5 - 214.5 °C; IR ν _{max} (Nujol)/cm ⁻¹ 3448, 3287, 3188, 3133,
8	D	83	3069, 2924, 2854, 1625, 1556 and 1468; NMR δ_H (400 MHz, DMSO)
			6.10 (2H, s), 6.64 – 6.67 (1H, m), 6.69 – 6.71 (1H, m), 7.09 (1H, dd, J
			3.5, 2.5 Hz), 7.21 – 7.24 (1H, d, J 2.5 Hz), 7.95 (1H, s), 11.22 (1H, s).
			IR v _{max} (Nujol)/cm ⁻¹ 3208, 3140, 2730, 1668, 1603, 1557, 1463, 1376
			and 1363; NMR δ_{H} (400 MHz, DMSO) 4.51 (2H, d, J 5.5 Hz), 6.68 –
9	F	19	6.70 (1H, m), 6.82 – 6.85 (1H, m), 7.07 (1H, d, J 3.5 Hz), 7.27 – 7.33
			(1H, m), 7.35 – 7.44 (5H, m), 7.95 (1H, d, J 1.0 Hz), 9.42 (1H, s), 9.86
			(1H, t, J 5.5 Hz), 11.88 (1H, s).
			mp 126.8 – 127.1 °C; IR v _{max} (Nujol)/cm ⁻¹ 2923, 2854, 1601, 1547,
			1458, 1376, 1256, 920 and 733; NMR $\delta_{\rm H}$ (400 MHz, DMSO) 5.52 (2H,
10	В	76	s), 6.80 – 6.83 (1H, m), 7.06 (1H, d, J 3.5 Hz), 7.10 – 7.19 (2H, m), 7.22
10			-7.28 (1H, m), 7.35 - 7.41 (1H. m), 7.52 (1H, d, J 3.5 Hz), 7.71 (1H, d,
			J 3.5 Hz), 8.10 (1H, s); Anal. Calcd for $C_{17}H_{11}ClN_3O + 0.25 H_2O$: C,
			61.46; H, 3.49, N, 12.65. Found: C, 61.31; H, 3.31; N, 12.37.
			mp 115.1 – 115.2 °C; NMR $\delta_{\rm H}$ (400 MHz, DMSO) 3.18 (6H, s), 5.35
11	C	86	(2H, s), 6.70 - 6.75 (2H, m), 7.10 - 7.27 (4H, m), 7.30 - 7.38 (2H, m),
			7.97 (1H, s); Anal. Calcd for $C_{19}H_{17}FN_4O + 0.1 H_2O$: C, 67.48; H, 5.13,
			N, 16.57. Found: C, 67.51; H, 5.10; N, 16.13; M/Z 336 M ⁺ .
		88	IR v _{max} (Nujol)/cm ⁻¹ 3252, 3112, 2924, 2854, 1582, 1515 and 1464;
			NMR $\delta_{\rm H}$ (400 MHz, DMSO) 3.67 (3H, s), 3.69 (3H, s), 4.46 (2H, d, J 6.5
12	_		Hz), 5.33 (2H, s), 6.69 – 6.75 (2H, m), 6.79 (1H, d, J 8.5 Hz), 6.89 (1H,
12		00	d, J 8.0 Hz), 6.97 – 7.09 (3H, m), 7.14 – 7.37 (5H, m), 7.96 (1H, s); Anal.
			Calcd for C ₂₆ H ₂₃ FN ₄ O ₃ : C, 68.11; H, 5.06, N, 12.21. Found: C, 68.14; H,
			5.08; N, 12.20.
	<u> </u>	<u> </u>	

			IR v_{max} (Nujol)/cm ⁻¹ 3329, 3193, 2924, 1649, 1577 and 1552; NMR δ_{H}
1.2	1		(400 MHz, DMSO) 5.34 (2H, s), 6.26 (2H, s), 6.70 – 6.73 (1H, m), 6.77
13 D	31	(1H, d, J 3.5 Hz), 6.95 (1H, t, J 8.0 Hz), 7.09 – 7.18 (2H, m), 7.19 – 7.28	
			(2H, m), 7.30 – 7.37 (1H, m), 7.96 (1H, s).

Adenosine Receptor Binding

5 Binding Affinities at hA_{2A} Receptors

The compounds were examined in an assay measuring *in vitro* binding to human adenosine A_{2A} receptors by determining the displacement of the adenosine A_{2A} receptor selective radioligand [3 H]-CGS 21680 using standard techniques. The results are summarised in Table 3.

10

Table 3

Example	K _i (nM)
Example 1	60
Example 8	272
Example 13	16

Evaluation of potential anti-Parkinsonian activity in vivo

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Haloperidol-induced hypolocomotion model

It has previously been demonstrated that adenosine antagonists, such as theophylline, can reverse the behavioural depressant effects of dopamine antagonists, such as haloperidol, in rodents (Mandhane S.N. et al., Adenosine A₂ receptors modulate haloperidol-induced catalepsy in rats. Eur. J. Pharmacol. 1997, 328, 135 - 141). This approach is also considered a valid method for screening drugs with potential antiparkinsonian effects. Thus, the ability of novel adenosine antagonists to block haloperidol-induced deficits in locomotor activity in mice can be used to assess both in vivo and potential antiparkinsonian efficacy.

Method

Female TO mice (25-30g) obtained from TUCK, UK, are used for all experiments. Animals are housed in groups of 8 [cage size – 40 (width) x 40 (length) x 20 (height)cm] under 12hr light/dark cycle (lights on 08:00hr), in a temperature ($20 \pm 2^{\circ}$ C) and humidity ($55 \pm 15\%$) controlled environment. Animals have free access to food and water, and are allowed at least 7 days to acclimatize after delivery before experimental use.

Drugs

Liquid injectable haloperidol (1 ml Serenance ampoules from Baker Norton, Harlow, 10 Essex, each containing haloperidol BP 5 mg, batch # P424) are diluted to a final concentration of 0.02 mg/ml using saline. Test compounds are typically prepared as aqueous suspensions in 8% Tween. All compounds are administered intraperitoneally in a volume of 10 ml/kg.

15 Procedure

1.5 hours before testing, mice are administered 0.2 mg/kg haloperidol, a dose that reduces baseline locomotor activity by at least 50%. Test substances are typically administered 5-60 minutes prior to testing. The animals are then placed individually into clean, clear polycarbonate cages [20 (width) x 40 (length) x 20 (height) cm, with a flat perforated, Perspex lid]. Horizontal locomotor activity is determined by placing the cages within a frame containing a 3 x 6 array of photocells linked to a computer, which tabulates beam breaks. Mice are left undisturbed to explore for 1 hour, and the number of beams breaks made during this period serves as a record of locomotor activity which is compared with data for control animals for statistically significant differences.

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6-OHDA Model

Parkinson's disease is a progressive neurodegenerative disorder characterised by symptoms of muscle rigidity, tremor, paucity of movement (hypokinesia), and postural instability. It has been established for some time that the primary deficit in PD is a loss of dopaminergic neurones in the substantia nigra which project to the striatum, and indeed a substantial proportion of striatal dopamine is lost (ca 80-85%) before symptoms are observed. The loss of striatal dopamine results in abnormal activity of the basal ganglia, a series of nuclei which regulate smooth and well co-ordinated movement (Blandini F. et al., Glutamate and Parkinson's Disease. *Mol. Neurobiol.* 1996, 12, 73 - 94). The neurochemical deficits seen

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in Parkinson's disease can be reproduced by local injection of the dopaminergic neurotoxin 6-hydroxydopamine into brain regions containing either the cell bodies or axonal fibres of the nigrostriatal neurones.

By unilaterally lesioning the nigrostriatal pathway on only one-side of the brain, a behavioural asymmetry in movement inhibition is observed. Although unilaterally-lesioned animals are still mobile and capable of self maintenance, the remaining dopamine-sensitive neurones on the lesioned side become supersenstive to stimulation. This is demonstrated by the observation that following systemic administration of dopamine agonists, such as apomorphine, animals show a pronounced rotation in a direction contralateral to the side of lesioning. The ability of compounds to induce contralateral rotations in 6-OHDA lesioned rats has proven to be a sensitive model to predict drug efficacy in the treatment of Parkinson's Disease.

15 Animals

Male Sprague-Dawley rats, obtained from Charles River, are used for all experiments. Animals are housed in groups of 5 under 12hr light/dark cycle (lights on 08:00hr), in a temperature (20 ± 2°C) and humidity (55 ± 15%) controlled environment. Animals have free access to food and water, and are allowed at least 7 days to acclimatize after delivery before experimental use.

Drugs

Ascorbic acid, desipramine, 6-OHDA and apomorphine (Sigma-Aldrich, Poole, UK). 6-OHDA is freshly prepared as a solution in 0.2% ascorbate at a concentration of 4 mg/mL 25 prior to surgery. Desipramine is dissolved in warm saline, and administered in a volume of 1 ml/kg. Apomorphine is dissolved in 0.02% ascorbate and administered in a volume of 2 mL/kg. Test compounds are suspended in 8%Tween and injected in a volume of 2 mL/kg.

Surgery

15 minutes prior to surgery, animals are given an intraperitoneal injection of the noradrenergic uptake inhibitor desipramine (25 mg/kg) to prevent damage to non-dopamine neurones. Animals are then placed in an anaesthetic chamber and anaesthetised using a mixture of oxygen and isoflurane. Once unconscious, the animals are transferred to a stereotaxic frame, where anaesthesia is maintained through a mask. The top of the animal's PCT/GB02/00095

head is shaved and sterilised using an iodine solution. Once dry, a 2 cm long incision is made along the midline of the scalp and the skin retracted and clipped back to expose the skull. A small hole is then drilled through the skill above the injection site. In order to lesion the nigrostriatal pathway, the injection cannula is slowly lowered to position above the right medial forebrain bundle at -3.2 mm anterior posterior, -1.5 mm medial lateral from bregma, and to a depth of 7.2 mm below the duramater. 2 minutes after lowing the cannula, 2 µL of 6-OHDA is infused at a rate of 0.5 µL/min over 4 minutes, yeilding a final dose of 8 µg. The cannula is then left in place for a further 5 minutes to facilitate diffusion before being slowly withdrawn. The skin is then sutured shut using Ethicon W501 Mersilk, and the animal removed from the strereotaxic frame and returned to its homecage. The rats are allowed 2 weeks to recover from surgery before behavioural testing.

Apparatus

Rotational behaviour is measured using an eight station rotameter system provided by Med Associates, San Diego, USA. Each station is comprised of a stainless steel bowl (45 cm diameter x 15 cm high) enclosed in a transparent Plexiglas cover running around the edge of the bowl, and extending to a height of 29 cm. To assess rotation, rats are placed in cloth jacket attached to a spring tether connected to optical rotameter positioned above the bowl, which assesses movement to the left or right either as partial (45°) or full (360°) rotations.

All eight stations are interfaced to a computer that tabulated data.

Procedure

To reduce stress during drug testing, rats are initially habituated to the apparatus for 15 minutes on four consecutive days. On the test day, rats are given an intraperitoneal injection of test compound 30 minutes prior to testing. Immediately prior to testing, animals are given a subcutaneous injection of a subthreshold dose of apomorphine, then placed in the harness and the number of rotations recorded for one hour. The total number of full contralatral rotations during the hour test period serves as an index of antiparkinsonian drug efficacy.

CLAIMS

1. Use of a compound of formula (I):

$$R_4$$
 R_5
 R_5
 R_1
 R_5
 R_1

5

wherein

R₁ is selected from alkyl, alkoxy, aryloxy, alkylthio, arylthio, aryl, halogen, CN, NR₇R₈, NR₆COR₇, NR₆CONR₇R₈, NR₆CO₂R₉ and NR₆SO₂R₉;

R₂ is selected from aryl attached via an unsaturated carbon;

10 R₃ and R₄ are independently selected from H, alkyl, halogen, alkoxy, alkylthio, CN and NR₇R₈;

R₅ is selected from H, acyclic alkyl, COR₆, CONR₇R₈, CONR₆NR₇R₈, CO₂R₉ and SO₂R₉; R₆, R₇ and R₈ are independently selected from H, alkyl and aryl or where R₇ and R₈ are in an NR₇R₈ group R₇ and R₈ may be linked to form a heterocyclic group, or where R₆, R₇ and R₈ are in a (CONR₆NR₇R₈) group, R₆ and R₇ may be linked to form a heterocyclic group; and R₉ is selected from alkyl and aryl,

or a pharmaceutically acceptable salt or prodrug thereof, in the manufacture of a medicament for the treatment or prevention of a disorder in which the blocking of purine receptors may be beneficial.

- 2. Use according to claim 1 wherein R₁ is selected from alkyl, alkoxy, thioalkyl, NR₇R₈, NR₆COR₇, NR₆CONR₇R₈, NR₆CO₂R₉ and NR₆SO₂R₉.
- 3. Use according to claim 1 wherein R₁ is selected from NR₇R₈, NR₆COR₇, NR₆CONR₇R₈, NR₆CO₂R₉ and NR₆SO₂R₉.
 - 4. Use according to claim 1 wherein R_1 is selected from NR_7R_8 .

ι,

- 5. Use according to claim 1 wherein R_1 is selected from NH_2 .
- 6. Use according to claim 1 wherein R₁ is selected from NR₆COR₇, NR₆CONR₇R₈, NR₆CO₂R₉ and NR₆SO₂R₉ and R₆ is H or alkyl.
 - 7. Use according to claim 1 wherein R₁ is selected from NR₆COR₇, NR₆CONR₇R₈, NR₆CO₂R₉ and NR₆SO₂R₉ and R₆ is H.
- 10 8. Use according to claim 1 wherein R_1 is selected from haloalkyl and arylalkyl.
 - 9. Use according to any preceding claim wherein R₂ is a 5- or 6- membered monocyclic aryl group.
- 15 10. Use according to any of claims 1 to 9 wherein R₂ is unsubstituted in at least one ortho position.
 - 11. Use according to any of claims 1 to 9 wherein R₂ is unsubstituted at both ortho positions.
 - 12. Use according to any preceding claim wherein R₂ is a heteroaryl group.
- 13. Use according to claim 12 wherein R₂ is a heteroaryl group which is attached to the pyrimidine ring of formula (I) such that a heteroatom is adjacent to said unsaturated carbon atom attached to the pyrimidine ring.
 - 14. Use according to any preceding claim wherein R₂ is an N, O or S-containing heteroaryl group.
- 30 15. Use according to any preceding claim wherein R₂ is selected from furyl, thienyl, pyridyl, pyrazolyl and thiazolyl.

- 16. Use according to any preceding claim wherein R₂ is selected from 2-furyl, 2-thienyl, 2-thiazolyl, 3-pyrazolyl and 2-pyridyl.
- 17. Use according to any preceding claim wherein R₃ is hydrogen.

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- 18. Use according to any preceding claim wherein R₄ is hydrogen.
- 19. Use according to any of claims 1 to 18 wherein R₅ is selected from H and substituted acyclic alkyl.

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- 20. Use according to claim 19 wherein R₅ is acyclic alkyl substituted by aryl, cycloalkyl, non-aromatic heterocyclyl, CO₂R₆, CONR₇R₈, CONR₆NR₇R₈ and C(=NR₆)NR₇R₈.
- 21. Use according to claim 20 wherein R₅ is acyclic alkyl substituted by aryl or CONR₇R₈.

- 22. Use according to claim 21 wherein R₅ is methyl substituted by aryl or CONR₇R₈.
- 23. Use according to any of claims 1 to 18 wherein R₅ is selected from (CR₁₀R₁₁)_nR₁₂ wherein n is 1 to 6, R₁₀ and R₁₁ are independently selected from H, alkyl and aryl, and R₁₂ is selected from substituted and unsubstituted aryl, cycloalkyl, non-aromatic heterocyclic, CO₂R₆, CONR₇R₈, CONR₆NR₇R₈ and C(=NR₆)NR₇R₈.
 - 24. Use according to claim 23 wherein n is 1.
- 25 25. Use according to claim 23 or 24 wherein R₁₀ and R₁₁ are independently selected from H and alkyl, preferably H.
 - 26. Use according to claim 23, 24 or 25 wherein R_{12} is selected from aryl and CONR₇R₈.
- 30 27. Use according to any of claims 23 to 26 wherein R_{12} is selected from mono-, di- or trisubstituted aryl groups represented by the formula $Ar(R_{13})_a(R_{14})_b(R_{15})_c$ wherein Ar is an aryl group; wherein R_{13} , R_{14} and R_{15} are substituent group(s), the same or different; and wherein a, b and c are 0 or 1 such that $a+b+c \ge 1$.

- 28. Use according to claim 26 or 27 wherein said aryl group is selected from phenyl, thienyl, furyl, indolyl and pyridyl.
- 5 29. Use according to claim 27 wherein R₁₃, R₁₄ and R₁₅ are independently selected from NR₇R₈, alkyl, alkoxy, halogen, NO₂, CN, hydroxy, NHOH, CHO, CONR₇R₈, CO₂R₇, NR₆CO₂R₉, NR₆SO₂R₉, OCO₂R₉ and aryl.
- 30. Use according to claim 29 wherein R_{13} , R_{14} and R_{15} are independently selected from NR_7R_8 , alkyl and halogen.
 - 31. Use according to claim 29 wherein R_{13} , R_{14} and R_{15} are independently selected from alkyl, and said alkyl is substituted alkyl and is selected from alkoxyalkyl, hydroxyalkyl, aminoalkyl and haloalkyl.

- 32. Use according to claim 30 wherein R_{13} , R_{14} and R_{15} are independently selected from NH₂, unsubstituted alkyl and fluoro.
- 33. Use according to claim 23, 24 or 25 wherein R₁₂ is selected from CONR₇R₈, R₇ is H and R₈ is selected from H, unsubstituted alkyl and arylalkyl.
 - 34. Use according to any one of claims 1 to 18 wherein R₅ is selected from CONR₇R₈, R₇ is H and R₈ is selected from arylalkyl.
- 25 35. Use according to claim 34 wherein R₈ is selected from arylmethyl.
 - 36. Use according to any preceding claim wherein R_6 to R_{11} are independently selected from lower alkyl.
- 30 37. Use according to claim 1 wherein R_1 is NH_2 , R_2 is 2-furyl, R_3 and R_4 are H and R_5 is arylmethyl.

- 38. Use according to claim 1 wherein the compound of formula (I) is 7-(2-fluorobenzyl)-4-(2-furyl)-7H-pyrrolo[2,3-d]pyrimidine-2-amine.
- 39. A method of treating or preventing a disorder in which the blocking of purine receptors may be beneficial comprising administration to a subject in need of such treatment an effective dose of a compound as set out in any one of claims 1 to 38 or a pharmaceutically acceptable salt thereof.
- 40. A use or method according to any one of claims 1 to 39 wherein the disorder is caused by the hyperfunctioning of purine receptors.
 - 41. A use or method according to any one of claims 1 to 40 wherein the purine receptors are adenosine receptors.
- 15 42. A use or method according to claim 41 wherein the adenosine receptors are A_{2A} receptors.
 - 43. A use or method according to any preceding claim wherein the disorder is a movement disorder.

- 44. A use or method according to claim 43 wherein the movement disorder is Parkinson's disease.
- 45. A use or method according to claim 44 for treatment of drug-induced Parkinsonism, post-encephalitic Parkinsonism, Parkinsonism induced by poisoning or post-traumatic Parkinson's disease.
- 46. A use or method according to claim 43 wherein the movement disorder is progressive supernuclear palsy, Huntingtons disease, multiple system atrophy, corticobasal degeneration, Wilsons disease, Hallerrorden-Spatz disease, progressive pallidal atrophy, Dopa-responsive dystonia-Parkinsonism, spasticity or other disorders of the basal ganglia which result in dyskinesias.

47. A use or method according to any one of claims 43 to 46 wherein the compound of formula (I) is in combination with one or more additional drugs useful in the treatment of movement disorders, the components being in the same formulation or in separate formulations for administration simultaneously or sequentially.

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- 48. A use or method according to claim 47 wherein said additional drug(s) useful in the treatment of movement disorders is/are a drug useful in the treatment of Parkinson's disease.
- 10 49. A use or method according to claim 47 or 48 wherein the or one of the additional drugs is L-DOPA or a dopamine agonist.
 - 50. A use or method according to any one of claims 1 to 42 wherein said disorder is depression, memory or cognitive impairment, acute or chronic pain, ADHD or narcolepsy.

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- 51. A use or method according to claim 50 wherein said memory or cognitive impairment disorder is Alzheimer's disease.
- 52. Use of a compound as set out in any one of claims 1 to 38 or a pharmaceutically acceptable salt thereof in the manufacture of a medicament for neuroprotection in a subject.
 - 53. A method of neuroprotection comprising administration to a subject in need of such treatment an effective dose of a compound as set out in any one of claims 1 to 38 or a pharmaceutically acceptable salt thereof.

- 54. A use or method according to claim 52 or 53 wherein said medicament or said method is for neuroprotection in a subject suffering from or at risk from a neurodegenerative disorder.
- 30 55. A use or method according to claim 54 wherein said neurodegenerative disorder is a movement disorder.

- 56. A use or method according to claim 55 wherein said movement disorder is a disorder as set out in claim 44, 45 or 46.
- 57. A use or method according to any one of claims 1 to 56 wherein the subject is human.
 - 58. A compound according to any one of claims 1 to 38, or a pharmaceutically acceptable salt thereof, for use in therapy.
- 10 59. A compound of formula (I) according to any one of claims 1 to 38, or a pharmaceutically acceptable salt or prodrug thereof, *per se*, wherein R₂ is selected from heteroaryl, other than compounds wherein R₂ is selected from pyrrolopyrimidines.
- 60. A compound according to claim 59 wherein R₂ is selected from 5- or 6- membered monocyclic heteroaryl groups.

Relevant to claim No.

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 A61K31/505 C07D487/04 A61P25/28 //(C07D487/04,239:00, 209:00)

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Category °

C, DOCUMENTS CONSIDERED TO BE RELEVANT

Minimum documentation searched (classification system followed by classification symbols) IPC 7 CO7D A61K A61P

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, WPI Data, BEILSTEIN Data, CHEM ABS Data, BIOSIS

Citation of document, with indication, where appropriate, of the relevant passages

Α	WO 99 62518 A (CADUS PHARMACE); MCKIBBEN BRYAN (US); WITTER 9 December 1999 (1999-12-09) claim 1	
A	WO 99 21617 A (BARALDI PIER G; MEDCO RES INC (US)) 6 May 1999 (1999-05-06) claim 1	IOVANNI 1-60
A	WO 99 01439 A (DU PONT PHARM 14 January 1999 (1999-01-14) claim 1	CO) 1-60
[V] 5	ther documents are listed in the continuation of box C.	Patent family members are listed in annex.
"A" docume cons "E" earlier filling "L" docume whice citati "O" docume other	categories of cited documents: nent defining the general state of the art which is not sidered to be of particular relevance or document but published on or after the international date of the document but published on priority claim(s) or the cited to establish the publication date of another on or other special reason (as specified) ment referring to an oral disclosure, use, exhibition or means then published prior to the international filing date but than the priority date claimed	 "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family
Date of the	e actual completion of the international search	Date of mailing of the international search report
	25 April 2002	22/05/2002
Name and	I mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Bliswijk	Authorized officer

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C.(Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	•
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	COCUZZA A J ET AL: "Use of the suzuki reaction for the synthesis of aryl-substituted heterocycles as corticotropin-releasing hormone (CRH) antagonists" BIOORGANIC & MEDICINAL CHEMISTRY LETTERS, OXFORD, GB, vol. 9, no. 7, 5 April 1999 (1999-04-05), pages 1063-1066, XP004162586 ISSN: 0960-894X page 1064	1-60
X	CHEMICAL ABSTRACTS, vol. 118, no. 3, 18 January 1993 (1993-01-18) Columbus, Ohio, US; abstract no. 22077, SUZUKI, HITOMI ET AL: "Sodium telluride in N-methyl-2-pyrrolidone. Reduction of aromatic carbonyl compounds to alcohols and formation of pyrrolo'2,3-d!pyrimidines (7-deaza-9H-purines) from aromatic nitriles" XP002197327 RN:144587-22-4,144587-23-5,144587-24-6 abstract & J. ORG. CHEM. (1993), 58(1), 241-4,	59
X	CHEMICAL ABSTRACTS, vol. 121, no. 9, 29 August 1994 (1994-08-29) Columbus, Ohio, US; abstract no. 108677, BOUILLON, JEAN PHILIPPE ET AL: "Trifluoromethylated pyrimidines starting from.betatrifluoroacetyl- lactams, -lactone and -cyclanone" XP002197328 RN:156870-48-3 abstract & HETEROCYCLES (1994), 37(2), 915-32,	59

			PC1/GB 02/00095	
C.(Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	•		
Category °	Citation of document, with indication, where appropriate, of the relevant passages		Relevant to claim No.	
X	CHEMICAL ABSTRACTS, vol. 113, no. 7, 13 August 1990 (1990-08-13) Columbus, Ohio, US; abstract no. 59080, KONDO, YOSHINORI ET AL: "Condensed heteroaromatic ring systems. XVI. Synthesis of pyrrolo'2,3-d!pyrimidine derivatives" XP002197329 RN:128266 abstract & CHEM. PHARM. BULL. (1989), 37(11), 2933-6,		59	
	·			

Ī	PCT/GB	02/00095
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Patent document cited in search report		Publication date		Patent family member(s)	Publication date
WO 9962518	A	09-12-1999	AU	4226599 A ·	20-12-1999
			BR	9911612 A	06-02-2001
			CN	1311680 T	05-09-2001
			CZ	20004443 A3	13-03-2002
			EP	1082120 A1	14-03-2001
			NO	20006090 A	31-01-2001
			TR	200003513 T2	21-06-2001
			MO	9962518 A1	09-12-1999
	•		US	2002028782 A1	07-03-2002
WO 9921617	Α	06-05-1999	US	5939432 A	17-08-1999
			US	6323214 B1	27-11-2001
			AU	9671798 A	17-05-1999
			CA	2302396 A1	06-05-1999
			EP	1025106 A2	09-08-2000
			MO	9921617 A2	06-05-1999
			US	6177444 B1	23-01-2001
			US	6194449 B1	27-02-2001
			US	2001047008 A1	29-11-2001
WO 9901439	Α	14-01-1999	AU	8181098 A	25-01-1999
			EP	0994860 A1	26-04-2000
			WO	9901439 A1	14-01-1999
			US	6103737 A	15-08-2000